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Role of Tissue Plasminogen Activator (tPA) in COVID-19 Associated Respiratory Distress Syndrome: A Review of Published Case Reports

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Abstract

Background: Articles published during the COVID-19 pandemic strongly suggest the association between a hyperinflammatory state and consumptive coagulopathy pathology in critically ill COVID-19 patients. Accordingly, the use of fibrinolytic agents, in CO-VID-19 patients with acute respiratory distress syndrome (ARDS) would be an efficacious approach. While evidence remains limited, certain published case reports have demonstrated improvement in the PaO2/FiO2 ratio in critical patients on tPA. This warrants a literature review to understand the therapeutic effectiveness of tPA in critically ill COVID-19 patients.

Methodology: An extensive literature review on the use of tPA in critically ill COVID-19 patients using secondary sources identified 84 studies of which 42 were analyzed for further screening. Post this screening, we further narrowed our scope to a review of case reports and case series. Our analysis is based exclusively on a systematic review of 4 case reports and 5 case series concerning 24 critically ill patients showing signs and symptoms of COVID-19.

Result: Out of the 24 cases reviewed, the majority of the cases (22) developed ARDS out of which 20 patients underwent intubation followed by mechanical ventilation. The outcome was reported in terms of death (7 patients), criticality (1 patient), improvements in arterial blood gas analysis parameters (6 patients), complete clinical recovery (8 patients), and no outcome in 2 patients.

Conclusion: In this review, a majority of critically ill patients with COVID-19 showed favorable outcomes post-treatment with tPA. However, larger studies are needed to confirm these findings.

Keywords: COVID-19; SARS COV-2; Acute Respiratory Distress Syndrome; ARDS; Tissue Plasminogen Activator; tPA; rt-tPA

Introduction

The COVID-19 pandemic caused by the SARS COV-2 has impacted mankind on a global scale. It has led to a complete economic shutdown with the burden of mass morbidities and mortalities. The cause of deaths in COVID-19 patients has not been well established yet [1]. Few of the hypotheses that have been proposed pen-

dulate between adult respiratory distress syndrome (ARDS) and pulmonary micro-thrombosis mentioned as COVID induced coagulopathy (CIC) [2]. Autopsies on few deceased COVID-19 patients revealed the pulmonary endothelial injury and alveolar-capillary microthrombi [3]. A high rate of pulmonary thromboembolism has been seen in critically ill COVID 19 patients with marked elevations in D-dimer levels alone or with fibrinogen, and fibrinogen degradation products [4]. Multi-system organ dysfunction as a result of CIC has also been observed in some patients [2,5]. A literature review during the pandemic has demonstrated reduced mortality rates with anticoagulant treatment in critical patients [6]. Connecting the temporality, physicians at various centers have tried tissue plasminogen activator (tPA) is seriously ill patients with the clinical diagnosis and radiological evidence of pulmonary thrombosis. We systematically reviewed 9 articles to understand the usefulness of tPA in CIC and analyze the clinical characteristics and outcomes in critically ill patients.

Methods

We searched PubMed/Medline, Web of Science, SCOPUS, until 7th October 2020 for case reports and case series using these keywords: COVID-19, SARS-CoV-2, acute respiratory distress syndrome, ARDS, Hypoxia, Hypoxemia, Tissue plasminogen activator, tPA, rt-PA, thrombolytic. All the published case reports included in the final analysis were in English. Our search identified 84 studies in total. After removing the duplicates, there were 42 studies for further screening. After excluding the review and original articles and including only case reports and case series we found 9 articles comprised of 24 cases [1,2,7-13]. Data from the article were curated and summarized in the form of country of origin, age, and gender of the patients, their presenting complaint, any coexisting comorbidities, medical interventions during hospitalization, and their outcome. Continuous variables were presented as means ± standard deviations and categorical data as absolute values and percentages. All data extraction and descriptive analysis were performed using Microsoft Excel.

Results

A systematic analysis of 9 articles demonstrating the use of tPA in COVID patients who developed ARDS was carried out, of which 5 were written as case series and 4 were described as case reports. 24 patients who displayed diagnostic and clinical signs of ARDS in COVID positive patients were identified of the 24 cases reviewed, age ranging from 23 to 82 years (mean 52.8 years), 16 (67%) were males and 8 (33%) were females. None of the cases reported race. Most cases were reported from the United States 20 (83%) followed by India 3 (13%) and Greece 1 (4%).

The most common comorbidities reported included hypertension in 13 patients (54%), diabetes mellitus (37%) in 9 patients, hyperlipidemia in 6 patients (25%), morbid obesity in 6 patients (25%) (Table 1). The detailed list of comorbidities is mentioned in tables 1. COVID diagnosis was incidental in 6 (26%) patients; 1 ST-Elevation Myocardial Infarction(STEMI), 1 atrial fibrillation, 1 pneumopericardium, 1 neutropenia due to Acute Myeloid Leukemia (AML) management, 1 hematoma due to lupus anticoagulant, and 1 with buprenorphine withdrawal. The most common presenting symptoms were dyspnea in 13 patients (54%), fever in 10 (42%), cough in 8 (33%), generalized weakness or fatigue in 5 (21%), chest pain in 2 (8%), myalgia in 2 (8%), headache in 1 (4%), altered mental status in 1 (4%) and vomiting in 1 (4%) and signs of coagulopathy in 2 (8%). ARDS was present in 22 (92%) patients. Many patients reported more than one comorbidity and symptom. The time from the development of ARDS to the requirement of intubation in these patients varied from 1 to 14 days (median 3.5 days). Besides ARDS, acute kidney injury (AKI), as a COVID 19 complication, was present in 4 (17%) patients.

As outlined in table 2 different authors used different TPA doses and regimens. The most common dosage regimen used was 25mg over 2h followed by 25 mg continuous infusion over 22 hours in 10 (42 %) patients. Concomitant Heparin was administered in 19 patients (79%) out of which Low molecular weight heparin(LMWH) 11(46 %) and Unfractionated Heparin(UFH)8 (33%). Besides tape, the treatment (see table no. for the detailed regimes) included antimicrobial agents in addition to supportive therapy. Hydroxychloroquine (HCQ) was given in 10 patients (42%) overall; with azithromycin in 8 (33%) patients. Other agents used were penicillin or cephalosporin in 7 (29%) patients, piperacillin-tazobactam in 1 patient, and vancomycin in 1 patient. Immunosuppressive agents prescribed in these cases included tocilizumab in 2 patients and methylprednisolone in 2 patients. Convalescent plasma was administered in 2 patients. Mechanical support and vasopressors were needed in 20 (83%) and 6 (25%) patients respectively and the combination was required in patients.

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Author	Age (years), Sex (M/F)	Coun- try	Past medical history	Presenting symptoms	COVID induced Complica- tions	Mean time from ARDS to Intuba- tion (days)	Mechanical Sup- port	Treatment ad- ministered
Poor., et al.	55 F	USA	Diabetes Mel- litus, Morbid Obesity.	Not men- tioned	AKI	1	VCV	HCQ, ceftriaxone
Poor., et al.	62 F	USA	Diabetes Mel- litus, Morbid Obesity.	Not men- tioned	AKI	1	VCV	Not mentioned
Poor., et al.	58 M	USA	Diabetes Mel- litus, Morbid Obesity, HTN, COPD	Not men- tioned	AKI	1	VCV	Not mentioned
Poor., et al.	57 M	USA	Morbid Obe- sity, HTN	Not men- tioned	AKI	1	VCV	Convalascent Plasma
Chris- tie., <i>et</i> al.	72 M	USA	Hyperlipid- emia, HTN	Worsening SOB, Fever x7 days	Respiratory Failure	2	APRV-PCV(Day 18)	HCQ, azithromycin
Chris- tie., <i>et</i> al.	68 F	USA	HTN, Hyper- lipidemia, CVA, COPD	Worsening SOB, Altered mental status	Respiratory Failure	2	FiO2- 100%/50L/ min(Day3),70%/ 50L (Day 5)	Piperacllin tazo- bactum
Chris- tie., <i>et</i> al.	55 F	USA	HTN, Hyper- lipidemia, Type 2 DM, Asthama	Worsening SOB, Fever, Cough, Weak- ness x2.5 weeks	Respiratory Failure	3	NRB 100%	Azithromycin, aug- mentin, levoflox, HCQ, MPS
Chris- tie., <i>et</i> al.	78 F	USA	Type 2 DM, CHF, HTN,CKD, CVA	Dyspnea, Fever	Respiratory Failure	1	PCV-VG with a FiO2 of 100% and PEEP of 15cmH2O.	Not mentioned
Chris- tie., et al.	82 M	USA	HTN, Hyper- lipidemia	Dyspnea, Cough x1 week	Respiratory Failure	3	BiPAP with a FiO2 of 100%	Not mentioned
Wang	78 M	USA	HTN, Hyper- lipidemia, CAD	Cough, Fa- tigue, Fever x1 week	Atrial Fibril- lation and Hypotension	6	None	HCQ, Azithromycin
Wang	59 F	USA	HTN	Cough, Myal- gia, Headache x2 days	Buprenor- phine withdrawal, Physical stress due to influenza	4	Acute respiratory distress	HCQ, Azithromycin
Wang	49 M	USA	None	Cough, Wors- ening SOB, Fever x6 days	Pneumoperi- cardium	1	None	HCQ, Azithromycin

Kari-	68 M	USA	Lung Cancer,	SOB, Chest	Respiratory	Few days	CPR support	325 mg Aspirin,
yanna	00 11	0011	Asthma, STEMI	Pain x2 hours	Failure	i cw ddys		600 mg Plavix
Molina., <i>et al</i> .	23M	USA	Nitrous Oxide abuse, Lupus Anticoagu- lant, Left sided neck Hematoma	Diffuse Weak- ness B/L LL	Coagulopa- thy	No ARDS	Not required	Vancomycin, ce- fepime, vit B12
Pa- pami- chalis	68M	Greece	HTN, DM, AML, Neutro- penia	Fever x7 days	Respiratory Failure, Co- agulopathy	10	APRV PEEP 8	Clarithromycin and Oseltamivir, hydroxychloquine, azithromycin, broadspec- trum antibiotic (meropenem), Tocilizumab 400 mg,Convalascent Plasma Day 35
Barett., <i>et al</i> .	39M	USA	None	SOB, Chest Pain, Cough	Respiratory Failure	4	APRV PEEP16	ceftriaxone, azithromycin, hy- droxychloroquine, Tocilizumab
Barett., <i>et al</i> .	58 M	USA	HTN, NIDM	SOB x2 weeks	None	14	None	ceftriaxone, azithromycin
Barett., <i>et al</i> .	67 M	USA	HTN, Thyroid Cancer	SOB, Cough, Fatigue, Fever x10 days	АКІ	10	APRV FiO2 -100%	ampicillin/sulbac- tam, hydroxychlo- roquine,
Barett., <i>et al</i> .	27 M	USA	Morbid Obe- sity, NIDM	Cough, Fever, SOB x7days	None	7	APRV FiO2 -82%	Not mentioned
Barett., et al.	52M	USA	Aortic Valve Disease, Hy- perlipidemia, Hodgkins Lymphoma	SOB, Fatigue, Fever, Body- ache x4 days	None	4	Not mentioned	ceftriaxone, azithromycin, hy- droxychloroquine
Goyal., et al.	45M	India	DM	Worsening Dyspnea	None	Not appli- cable	FiO2-0.7	Low molecular weight heparin (LMWH)
Goyal., et al.	60F	India	CAD, HTN	SOB, Vomiting x4 days	None	Not appli- cable	FiO2-0.75	SPO2-52%
Goyal., <i>et al</i> .	59F	India	Hypothyroid- ism	SOB, Fever x2 days	None	Not appli- cable	FiO2-0.7	Not mentioned
Sethi., <i>et al</i> .	44M	USA	DM, Obesity	SOB, Cough	hypoxemic respiratory	Not appli- cable	FiO2-86%	methylpredniso- lone

Table 1: Demographics, Co-morbidities, Symptoms, Complications and Treatment in COVID Patients.

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Au- thor	TPA regimen	O2 require- ments pre TPA infu- sion	O2 require- ments post TPA	ABG findings pre TPA	ABG findings post TPA trans- fusion	Vaso- pressor require- ment pre TPA infusion	Vasopres- sor re- quirement Post TPA transfusion	D-dimer/Fibrin- ogen pre TPA transfusion	D-dimer/Fi- brinogen post TPA transfu- sion	Outcome
Poor., et al.	50mg over 2h fd by 2mg/h for 24h	FiO2-60% PEEP-15	Not men- tioned	pH -7.12/ PaC02-71/ PaO2-45	pH 7.27/PaCO2 40 mmHg, PaO2 78 mmHg.	norepi- nephrine 30 mcg/ min and vasopres- sin 2.4 units/h	norepineph- rine 4 mcg/ min	5.7μg/mL	Not mentioned	Death d/t Septic Shock
Poor., et al.	50mg over 2h	FiO2-70% PEEP-15	Not men- tioned	pH 7.33, PaCO2 55 mm Hg, and PaO2 115 mm Hg	pH 7.33, PaCO2 55 mm Hg, and PaO2 115 mm Hg	norepi- nephrine 15 mcg/ min	Not re- quired	6.1µg/mL	Not mentioned	Recovery 10 days
Poor., et al.	50mg over 75min	FiO2- 60%PEEP-5	Not men- tioned	pH 7.14, PaCO2 107 mm Hg and PaO2 84 mm Hg	pH 7.18, PaCO2 89 mm Hg, and PaO2 66 mm Hg,	norepi- nephrine 50 mcg/ min and vasopres- sin 2.4 units/h.	norepineph- rine dose was weaned to 7 mcg/ min	4.6 μg/mL	Not mentioned	Death / ECHO s/o biven- tricular thrombi
Poor., et al.	50mg over 2h fd by 2mg/h	FiO2-100% PEEP-16	Not men- tioned	pH 7.21, PaCO2 51 mm Hg, PO2 81 mm Hg	pH 7.27, PaCO2 51 mm Hg, and PaO2 140 mm Hg	norepi- nephrine 10 mcg/ min	Not men- tioned	6.6 μg/mL	Not mentioned	Death d/t Shock
Chris- tie., et al.	25mg over 2h fd by 25 mg con- tinuous infusion over 22 hours	Day 1 50% venturi mask, Day 2 high flow nasal cannula (HFNC) 60%/40L/ min, 100%/60L/ min	FiO2 of 80% (Day 11)	PaO2 53 mm Hg, PaO2/ FiO2 (P/F) ratio ranging from 69 (Day 10)	P/F ratio increased to 76 (Day 11), P/F ratio -121 (Day 15), P/F ratio -127	Not re- quired	Not re- quired	D -dimer- 2.16ug/ mL and Fibrino- gen -654mg/dL	D-dimer to 9.57ug/mL (Day 11) D dimer 1.99ug/ mL.	Recovery Day 18
Chris- tie., et al.	25mg over 2h fd by 25 mg con- tinuous infusion over 22 hours	100% non- rebreather (NRB), HFNC 60%/30L/ min (Day 2), 100%/70L/ min (Day 3)	45%40L/ min (Day 4)	PaO2-72 mm Hg,	Sp02 increased from 71% to 89%	Not re- quired	Not re- quired	D-dimer fibrino- gen were elevated at 1.87ug/mL and 512mg/dL	D-dimer ini- tially increased to 5.57ug/ mL and her fibrinogen decreased to 475mg/dL, D-dimer had decreased to 2.39ug/mL (Day 5)	Aspiration Pneumo- nia

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Chris- tie., et al.	25mg over 2h fd by 25 mg con- tinuous infusion over 22 hours	NRB 100% (Day 4)	NRB 100% (Day 6),6L Nasal Can- nula (Day 8)	PaO2 of 51mmHg (Day 1), PaO2 of 67mmHg (Day 4), PaO2 59mmHg (Day 5)	PaO2 72mmHg (Day 6), PaO2 77mmHg (Day 7)	Not re- quired	Not re- quired	D-dimer of 8.34ug/mL and fibrinogen of 899mg/dL	D-dimer increased to >20ug/ mL, and her fibrinogen decreased to 535mg/dL (Day 6),D- dimer stayed above 20ug/m (Day 7),r D-di- mer decreased to 4.56ug/ mL (Day 8), D-dimer at this point was 1.91ug/mL (Day 11)	Recovery Day 13
Chris- tie., et al.	25mg over 2h fd by 25 mg con- tinuous infusion over 22 hours	NRB 100%/15L/ min (Day 1), HFNC 100%/70L/ min (Day 3)	FiO2 require- ment was decreased from 100% to 45%	PaO2 of 48mmHg and SpO2 85% (Day 1), PaO2 of 61mmHg (Day 3)	P/F ratios ranged from 175 to 196 (Day 4), P/F ratios in the 190s (Day 5)	Required	Weaned off	D-dimer and fibrinogen were 2.47ug/mL and 744mg/dL (Day 3)	D-dimer increased to 7.05ug/ mL and her fibrinogen decreased to 596mg/dL	Recovery Day 10
Chris- tie., et al.	25mg over 2h fd by 25 mg con- tinuous infusion over 22 hours	NRB 100% (Day 1), HFNC 100%/50L/ min (Day 2), HFNC 100%/50L/ min (Day 3)	HFNC 100%60L/ min (Day 4)	PaO2 of 55mmHg and a SpO2 of 92% (Day 1), PaO2 of 57mmHg (Day 2), PaO2 of 67 mm Hg (Day 2)	PaO2 of 57mmHg (Day 3), PaO2 in- creased to 79mmHg (Day 3)	Not re- quired	Not re- quired	D-dimer and fibrinogen were 4.78ug/mL and 753mg/dL	D-dimer had increased to >20ug/mL and her fibrinogen had decreased to 693mg/dL (Day 3), D-di- mer decreased to 14.06ug/mL (Day4)	Recovery Day 4
Wang., et al.	25mg over 2h fd by 25mg over 22h	oxygen require- ment 4 to 6 L/min (Day 1), 100% fraction of inspired oxygen (FiO2) on a non- rebreather mask (NRB) by day 3	Fi02 >60%	(P/F) ratio was 73 (Day 6), P/F ratio between 140 and 240 (Day8)	P/F ratio had improved to 408 (Day 9). P/f ratio worsened to 136 (1 Hour post heparin infusion), P/F ratio- 188-250 (Day 10)	norepi- nephrine	Norepi- nephrine, Vasopressin and Phenyl- ephrine	D-dimer >50 000 ng/mL and fibrinogen levels ranged be- tween 375 and 541 mg/dL (Day 6-10)	Fibrinogen lev- els remained similar at 351 mg/dL and his D-dimer had decreased to 16 678 ng/	Death (Day 11)

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Wang.,	25mg	100% NRB	100%NRB	P/F ratio 82	P/F was 135 (4	None	Reduced	r D-dimer was	D-dimer in-	Improve-
et al.	over 2h fd by 25mg over 22h	(Day 2)		(Day 4)	hours post trans- fusion) P/F ratio had im- proved to 150 (12 hours post transfusion), P/F ratio was now 135 (38 hours post transfusion)	None		545 ng/mL, increased to 20 293 ng/mL by HD 9 with a fibrino- gen level of 939 mg/dL	creased to 40 490 ng/ mL (12 hours post transfu- sion)	ment in P/F ratio
Wang., et al.	25mg over 2h fd by 25mg over 22h	100% FiO2 via NRB (Day 1)	100%FiO2	P/F ratio was 120 (prone Day 1), P/F ratio ranged from 72 to 90 (supine Day 1)	P/F of 125 by 3 hours after completion of tPA,s P/F ratio declined to 71 (supine 33 hours post Tpa), P/F ratio of 118 (prone)	Required	NA	D-dimer was 33 228 ng/ mL (Day 1) it had reduced to 17 301 ng/ml (Day 2)	D-dimer 37 215 ng/mL and his fibrin- ogen 544 mg/ dL (35 hours post-tPA)	Improve- ment in P/F ratio
Kari- yanna., <i>et al</i> .	100 mg of Tpa	NA	CPR Re- quired	PaO2 - 68.6 ph-7.29 PaCo2 -18.1 (Day 1)	PaO2-30.7 ph- 7.35 PaCo2-17.5 (Day 2)	Not re- quired	ACLS			Death (Day 2)
Mo- lina., <i>et al</i> .	Not men- tioned	SPO2-92%	None	PCO2 of 37 mmHg and a venous pH of 7.4	Not mentioned	Not re- quired	Not re- quired	d-dimer of 7386 ng/mL (Day 1)	Not mentioned	Not men- tioned
Pa- pami- chalis	25mg over 2h fd by 25mg over 22h	Not men- tioned	Not men- tioned	(P/F) ratio 115 mmHg (Day 1), P/F ratio 241 mm Hg (Day 3-`10) P/F ratio 153 mmHg (Day 13)	P/F ratio 228 mmHg 15 hours after rt-PA initiation, P/F ratio <150 mm Hg 48 hours post infusion	Not re- quired	Not re- quired	D dimer 1.9 ng/ml (Day 1), D dimer -2.1 ng/ml (Day 3) D dimer -2.8 ng/ml (Day 6), D dimer -2.6 ng/ml (Day 8), D dimer -4.6 (Day 10), D dimer -4.9 ng/ml (Day 12)	D dimer -9.6 ng/mL (Day 13),D dimer -2.6 ng/mL (Day 35),D dimer -2.7 ng/ ml (Day 45)	Death Day 45
Bar- ett., <i>et</i> al.	25mg over 2h fd by 25mg over 22h,50 mg over 2 hrs		Not men- tioned	(P/F) ratio 81 (Day 5 supine position). P/F ratio 110 (Day 5 prone position), P/F ratio 60-100 (Day 8)	P/F ratio-197 (24 hours post tpa infusion), P/F ratio - 227 (36 hours post tpa infusion)	Not re- quired	Not re- quired	r fibrinogen of 1,116 mg/dL, D-dimer of 7,434 ng/mL (Day 5), Fibrinogen -731mg/dl	Fibrinogen -6	28mg/dl
Bar- ett., <i>et</i> <i>al</i> .	50 mg over 2 hrs x2	100%FiO2 via NRB (Day 1),	Not men- tioned	h P/F ratios persistently in the 90's (Day 5)	P/F ratio-136 (24 hours post tpa infusion), P/F ratio post second bolus - 90, P/F ratio -114,175	Not re- quired	Not re- quired	fibrinogen of 482 mg/dL (Day 1), D-dimer of 1,462 ng/mL (Day 1), pre-t PA fibrino- gen was 980 mg/ dL D-dimer 2,124 ng/mL (Day 5)	, post-tPA fibrinogen 944 mg/dL , Ddimer 7,094 ng/mL	Improve- ment in P/F ratio

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Bar- ett. <i>, et</i> al.	50 mg over 2 hrs x3	Spo2-80% (on admis- sion), FiO2 -80%	Fi02 -50%	P/F ratios ranging from 70-105, (pH 7.1-7.2) (Day 6), P/F ratio -77 (Day 16)	P/F ratio -92 (4 hours post tpa infusion), P/F ratio -85 (24 hours post tpa infusion), P/F ra- tio -105 (3 hours post second infusion)	Required	Required	fibrinogen 257 mg/dL, D-dimer 6,070 ng/mL (on admission), D dimer>35,000ng/ mL (Day 6)	Not mentioned	Death (Day 18)
Bar- ett., <i>et</i> al.	50 mg over 2 hrs, 25mg over 22h,	SpO2-80% on FiO2- 100% NRB (on admis- sion)	FiO2- 100% PEEP-15	P/F ratios in the 60's (prone posi- tioning),	P/F ratio was 217 (5 hours post tpa infu- sion), P/F ratio - 71 (completion of infusion)	Not men- tioned	Not men- tioned	fibrinogen 750 mg/dL, D-dimer 2,240 ng/Ml (Day 1), (peak 856 mg/dL on HD 2 and 4), pre-tPA fibrinogen was 756mg/dL and D- dimer was 4,040 ng/Ml	post-tPA fibrinogen 856 mg/dL D-dimer to >20,000 ng/ mL	Critical
Bar- ett., <i>et</i> <i>al.</i>	50 mg over 2 hrs	Sp02-82% 100% Fi02- 100% NRB (on admis- sion)	Not men- tioned	P/F ratio of 97 (supine Day 6), P/F ratio to >100 (prone Day 6) P/F <100 (prone Day 12)	P/F ratio imme- diately improved from 82 pre-tPA to 105 post-tPA (Day 12), P/F ra- tio had improved to 141 (Day 13)	Not re- quired	Not re- quired	fibrinogen of 836 mg/dL (peak 1,070 mg/dL on HD4), D-dimer of 843 ng/mL (Day 1), pre-tPA fibrinogen was 365 mg/dL and D-dimer 15,061 ng/mL (Day 12)	post-tPA fibrinogen was 373 mg/ dL and D- dimer17,613 ng/mL (Day 12)	Improve- ment in P/F ratio
Goyal., <i>et al</i> .	2mg/ hr,30mg over 15hr	FiO2-0.7 (NIV)	FiO2-0.5, FiO2-0.35 (Day 6)	Not men- tioned	Not mentioned	Not re- quired	Not re- quired	D dimer was 1350 ng/ml and fbrinogen was 670 mg/dl (Day 2)	Not mentioned	Recovery (Day 10)
Goyal., et al.	50 mg over 3 hr	FiO2-0.75	FiO2-0.6 (Day 2), FiO2-0.4 (Day 3), FiO2-0.21 (Day 7)	P/F ratio -90 (on admis- sion)	Not mentioned	Not re- quired	Not re- quired	D-dimer and fbrinogen were 1787 ng/ml and 704 mg/dl (Day 1)	Not mentioned	Recovery (Day 13)
Goyal., <i>et al</i> .	50 mg over 3 hr	SPO2-58% on room air, FiO2- 0.7HFNC	FiO2-0.35 (3 hours post trans- fusion)	P/F ratio was 80	FiO2-0.35	Not re- quired	Not re- quired	D-dimer and fbrinogen were 4583 ng/ml and 415 mg/ dl	Not mentioned	Recovery (Day 8)
Sethi., et al.	100 mg over 2h	FiO2-86%	Not men- tioned	pH -7.14/ PaC02-53/ PaO2-193	Not available	norepi- nephrine 40 mcg/ min and vasopres- sin 2.4 units/h	Pressors support weaned off.	>20 μg/mL	Not mentioned	Improved, r e m a i n hospital- ized

Table 2: TPA Regimen, Pre and Post TPA Parameters and Outcomes.

The following parameters-Arterial blood gases (ABG), oxygen requirements, vasopressor requirements, D dimer levels, and Fibrinogen degradation products (FDP) levels were reported pre and post tPA transfusion. A clinical reduction in hypoxia was seen as an improvement in P/F ratio in 5 (21%) patients; increase in PaO2 in 15(62 %) patients and improvement in D dimer and FDP levels 7(29%) and 5 (21%) patients respectively and both D dimer and FDP in 3 (13%) patients. The tPA dosing regimens were variable in the 9 articles that were reviewed. With the usage of 25mg over 2h followed by 25mg over 22h being the most commonly administered in 9 patients (38%) and 50 mg bolus in 8(33%) patients. As per the treatment outcomes reported, 8 (33%) patients showed complete clinical recovery, 6 (25%) patients reported improvement in P/F ratio, 1 (4%) patient was reported as critical and7 (29%) patients died, no outcome was reported for 2 patients.

Discussion

Autopsy studies from the Severe Acute Respiratory Syndrome (SARS) outbreak of the early 2000s caused by SARS COV-1 reported pulmonary thrombi [4,14,15]. SARS COV- 2 appears to show a similar pro-coagulable state causing gas exchange derangements and multi-organ dysfunction [1]. As stated by Goyal., *et al.* bleeding tendencies are rare in COVID 19 patients and coagulation pathway derangement are associated with a prothrombotic state [1]. Wright., *et al.* supported this hypothesis by showing an elevated D dimer level and complete failure of clot lysis at 30 minutes on thromboelastography in seriously ill COVID positive patients [16]. To counter this hypercoagulable state, tPA has been one of the treatment modalities used in critically ill COVID 19 patients.

To the best of our knowledge, this is the first review summarizing the evidence of the use of tPA in critically ill COVID 19 patients. In addition to tPA usage across studies, concomitant heparin was also prescribed in a majority of cases. Thrombolysis with tPA showed favorable clinical outcomes reported as improvements in PaO2, P/F ratio, and coagulation parameters. The use of tPA was associated with clinical improvement in 14 out of the 24 patients included in our analysis, 8 of which were described to have a clinical recovery, 6 showed improvements in oxygenation parameters. In COVID 19 induced ARDS, mortality rates reported in critically ill patients were 30-50% on High Flow Nasal Cannula (HFNC) to 80 to 90% in patients on mechanical ventilation [11]. Of the 9 studies that we included in our analysis, 8 studies considered the intervention of intubation with mechanical ventilation and the median time to intubation from ARDS was 3.5 days. The article by Goyal., et al. mentioned the preference of not intubate patients and using

thrombolysis as an alternative [1].

Some literature suggests that there seems to be a significant role of CIC in the pathogenesis of COVID-19 induced respiratory failure [17,18], although not all autopsy studies describe similar findings [19]. If this holds, rtPA may be an efficacious drug. However, Meyer, *et al.* in their study on "Fibrinolysis for Patients with Intermediate-Risk Pulmonary Embolism" reported an increased risk of major hemorrhage and stroke [20]. Therefore, given its recognized risks in patients with lower mortality risk, the benefit to risk ratio should be carefully considered.

Limitations

There are some inherent limitations to the current review; namely, a variability in baseline characteristics and demographics of patients included in the individual case reports/series. It is difficult to generalize the outcomes from a small set of patients to a larger population owing to the limited sample size of the analyzed data and variation in tPA dosing regimens used across studies. Furthermore, demographics and baseline information cannot be used for outcomes of a larger population without a control group. Besides, inconsistency in reporting information, making it difficult to generalize these outcomes to a larger population. Furthermore, clinical case reports have a high risk of publication bias as it is expected that only positive results will be published. Besides, heterogeneity of the included studies cannot be ignored. Small sample size could hinder a robust case selection which could eventually influence the benefit to risk ratio assessment.

Conclusion

The current review of the case reports analyzed the available literature on tPA use in severely ill COVID 19 patients. The use of tPA seems to have a favorable outcome in a noticeable number of cases. However, an analysis of clinical reports cannot replace evidence provided by clinical trials. Therefore, large studies are further needed on the topic to provide robust evidence to determine the effectiveness and optimal dosing regimen.

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